=> d his

(FILE 'HOME' ENTERED AT 05:43:37 ON 25 MAR 2002)

FILE 'REGISTRY' ENTERED AT 05:43:49 ON 25 MAR 2002

L1 STRUCTURE UPLOADED

L2 12 S L1

185 S L1 FULL L3

FILE 'HCAPLUS' ENTERED AT 05:45:00 ON 25 MAR 2002

L416 S L3

L5 8 S L3/THU

0 S L5 AND BULLINGTON, J?/AU L6

FILE 'CAOLD' ENTERED AT 05:49:51 ON 25 MAR 2002

=> s 13

L7 0 L3

STN Structure : 9580882c.str

```
13 18 19 20 21
ring nodes:
    1 2 3 4 5 6 7 8 9 10 11 12
ring/chain nodes:
    14
chain bonds:
    1-10 8-13 11-18 13-14 14-21 18-19 18-20
ring bonds:
    1-2 1-6 2-3 3-4 4-5 5-6 7-8 7-12 8-9 9-10 10-11 11-12
exact/norm bonds:
    7-8 7-12 8-9 9-10 10-11 11-12 14-21 18-19 18-20
exact bonds:
    1-10 8-13 11-18 13-14
normalized bonds:
    1-2 1-6 2-3 3-4 4-5 5-6
isolated ring systems:
    containing 1:
```

G1:0,5

chain nodes :

Match level:
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:CLASS 14:CLASS 18:CLASS 19:CLASS 20:CLASS 21:CLASS

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Trying 3106016892...Open
Welcome to STN International! Enter x:x
LOGINID:ssspta1612BXR
PASSWORD:
TERMINAL (ENTER 1, 2, 3, OR ?):2
                     Welcome to STN International
NEWS
                 Web Page URLs for STN Seminar Schedule - N. America
NEWS
                 IMSworld Pharmaceutical Company Directory name change
                 to PHARMASEARCH
NEWS
     3 Oct 09
                 Korean abstracts now included in Derwent World Patents
                 Index
NEWS 4 Oct 09
                 Number of Derwent World Patents Index updates increased
NEWS 5 Oct 15
                 Calculated properties now in the REGISTRY/ZREGISTRY File
NEWS 6 Oct 22 Over 1 million reactions added to CASREACT
NEWS 7 Oct 22 DGENE GETSIM has been improved
NEWS 8 Oct 29 AAASD no longer available
NEWS 9 Nov 19 New Search Capabilities USPATFULL and USPAT2
NEWS 10 Nov 19 TOXCENTER(SM) - new toxicology file now available on STN
NEWS 11 Nov 29 COPPERLIT now available on STN
NEWS 12 Nov 29 DWPI revisions to NTIS and US Provisional Numbers
NEWS 13 Nov 30 Files VETU and VETB to have open access
NEWS 14 Dec 10 WPINDEX/WPIDS/WPIX New and Revised Manual Codes for 2002
NEWS 15 Dec 10 DGENE BLAST Homology Search
NEWS 16 Dec 17 WELDASEARCH now available on STN
NEWS 17 Dec 17 STANDARDS now available on STN
NEWS 18 Dec 17 New fields for DPCI
NEWS 19 Dec 19 CAS Roles modified
NEWS 20 Dec 19 1907-1946 data and page images added to CA and CAplus
NEWS 21
         Jan 25 BLAST(R) searching in REGISTRY available in STN on the Web
NEWS 22 Jan 25
                 Searching with the P indicator for Preparations
NEWS 23
         Jan 29
                 FSTA has been reloaded and moves to weekly updates
NEWS 24 Feb 01
                 DKILIT now produced by FIZ Karlsruhe and has a new update
                 frequency
NEWS 25
         Feb 19
                 Access via Tymnet and SprintNet Eliminated Effective 3/31/02
NEWS 26 Mar 08
                 Gene Names now available in BIOSIS
         Mar 22
NEWS 27
                 TOXLIT no longer available
NEWS 28 Mar 22 TRCTHERMO no longer available
              February 1 CURRENT WINDOWS VERSION IS V6.0d,
NEWS EXPRESS
              CURRENT MACINTOSH VERSION IS V6.0a(ENG) AND V6.0Ja(JP),
              AND CURRENT DISCOVER FILE IS DATED 05 FEBRUARY 2002
              STN Operating Hours Plus Help Desk Availability
NEWS HOURS
NEWS INTER
              General Internet Information
NEWS LOGIN
              Welcome Banner and News Items
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FILE 'HOME' ENTERED AT 05:43:37 ON 25 MAR 2002

=> file req

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.15 0.15

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 05:43:49 ON 25 MAR 2002 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2002 American Chemical Society (ACS)

STRUCTURE FILE UPDATES: 22 MAR 2002 HIGHEST RN 402712-52-1 DICTIONARY FILE UPDATES: 22 MAR 2002 HIGHEST RN 402712-52-1

TSCA INFORMATION NOW CURRENT THROUGH July 7, 2001

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Calculated physical property data is now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

The P indicator for Preparations was not generated for all of the CAS Registry Numbers that were added to the H/Z/CA/CAplus files between 12/27/01 and 1/23/02. Use of the P indicator in online and SDI searches during this period, either directly appended to a CAS Registry Number or by qualifying an L-number with /P, may have yielded incomplete results. As of 1/23/02, the situation has been resolved. Also, note that searches conducted using the PREP role indicator were not affected.

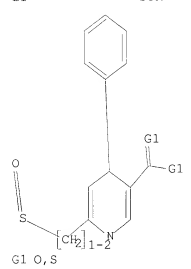
Customers running searches and/or SDIs in the H/Z/CA/CAplus files incorporating CAS Registry Numbers with the P indicator between 12/27/01 and 1/23/02, are encouraged to re-run these strategies. Contact the CAS Help Desk at 1-800-848-6533 in North America or 1-614-447-3698, worldwide, or send an e-mail to help@cas.org for further assistance or to receive a credit for any duplicate searches.

Uploading 9580882c.str

L1 STRUCTURE UPLOADED

=> d 11

L1 HAS NO ANSWERS L1 STR



Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 05:44:28 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 23 TO ITERATE

100.0% PROCESSED 23 ITERATIONS 12 ANSWERS SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**

PROJECTED ITERATIONS: 173 TO 747 PROJECTED ANSWERS: 33 TO 447

L2 12 SEA SSS SAM L1

=> s l1 full

THE ESTIMATED SEARCH COST FOR FILE 'REGISTRY' IS 139.90 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N or END:y

FULL SEARCH INITIATED 05:44:41 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 326 TO ITERATE

100.0% PROCESSED 326 ITERATIONS 185 ANSWERS

SEARCH TIME: 00.00.02

L3 185 SEA SSS FUL L1

=> file hcaplus

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 140.54 140.69

FULL ESTIMATED COST

FILE 'HCAPLUS' ENTERED AT 05:45:00 ON 25 MAR 2002 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2002 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 25 Mar 2002 VOL 136 ISS 13 FILE LAST UPDATED: 22 Mar 2002 (20020322/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

CAS roles have been modified effective December 16, 2001. Please check your SDI profiles to see if they need to be revised. For information on CAS roles, enter HELP ROLES at an arrow prompt or use the CAS Roles thesaurus (/RL field) in this file.

The P indicator for Preparations was not generated for all of the CAS Registry Numbers that were added to the CAS files between 12/27/01 and 1/23/02. As of 1/23/02, the situation has been resolved. Searches and/or SDIs in the H/Z/CA/CAplus files incorporating CAS Registry Numbers with the P indicator executed between 12/27/01 and 1/23/02 may be incomplete. See the NEWS message on this topic for more information.

=> s 13

L4 16 L3

=> s 14/thu

FIELD CODES CANNOT BE CHANGED HERE
You may have tried to apply a field code to a term that already has a
field code. You can only add a field code to a term that has no field
code appended to it.

=> s 13/thu

16 L3 428947 THU/RL L5 8 L3/THU

(L3 (L) THU/RL)

 \Rightarrow s 15 and bullington, j?/au

40 BULLINGTON, J?/AU

L6 0 L5 AND BULLINGTON, J?/AU

=> d 15, ibib abs fhitstr, 1-8

L5 ANSWER 1 OF 8 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1999:250272 HCAPLUS

DOCUMENT NUMBER: 130:332887

TITLE: Optically active 1,4-dihydropyridines as bradykinin

antagonists, their intermediates, preparation of

their

intermediates, and pharmaceutical compositions

containing them

INVENTOR(S): Ikeda, Takafumi; Kawamura, Mitsuhiro; Katsura,

Gokutei

PATENT ASSIGNEE(S): Pfizer Pharmaceutical Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 51 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND	DATE	APPLICATION NO. DATE
JP 11106375	A2	19990420	JP 1998-218686 19980717
EP 899261	A1	19990303	EP 1998-306202 19980804
R: AT, BE	, CH, DE,	DK, ES,	FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI	, LT, LV,	FI, RO	
US 6156752	А	20001205	US 1998-133580 19980813
CA 2245041		19990218	CA 1998-2245041 19980814
BR 9803180	А	20000321	BR 1998-3180 19980818
PRIORITY APPLN. INF			WO 1997-IB1000 A 19970818
OTHER SOURCE(S):	MAF	RPAT 130:3	32887
GI			

AB Bradykinin antagonists, 1,4-dihydroxypyridines I [A1, A2 = halo; R1, R2 = C1-4 alkyl; R3 = (substituted) Ph, naphthyl; Y = (substituted) heteocyclyl

selected from C5-10 azacycloalkyl, C6-10 diazacycloalkyl, and C7-10 azabicycloalkyl; R4 = (substituted) C1-8 alkyl, (substituted) amino, (substituted) C2-6 alkanoyl, (substituted) C3-8 cycloalkyl, (substituted) C7-14 bicycloalkyl, (substituted) C5-10 azacycloalkyl, (substituted) C6-10

diazacycloalkyl, (substituted) C7-14 monoazabicycloalkyl, (substituted) C7-14 diazabicycloalkyl] or their pharmaceutically acceptable salts are useful for pharmaceutical compns. contg. pharmaceutically acceptable carriers, for treatment of inflammation, cardiovascular diseases, pain, cold, allergy, asthma, pancreatitis, burn, viral infection, head injury, Alzheimer's disease, or multiple trauma. 1,4-Dihydropyridines II [A1, A2 = halo; R1, R2 = C1-4 alkyl; R3 = (substituted) Ph, (substituted) naphthyl; B1B2B3NH+ = chiral amine residue] are prepd. by oxidn. of III (A1, A2, R1-R3 = same as above; Z = H, C1-6 alkyl) with oxidizing agents and fractional crystn. of the resulting S-oxides using chiral amines. 1,4-Dihydroxypyridines IV (A1, A2, R1-R3 = same as above; R = C1-4 alkyl) are prepd. by chiral oxidn. of diastereoselective oxidn. of III in the presence of oxidizing agents. Di-Me

(-) - (4S) - 4 - (2, 6 - dichlorophenyl) - 2 - [4 -

(8-methyl-8-azabicyclo[3.2.1]oct-3-yl)-1-piperazinyl]carbonylmethyl-6-(S)-phenylsulfinylmethyl-1,4-dihydropyridine-3,5-dicarboxylate dicitrate, which inhibited the binding of bradykinin to B2 receptors in IMR-90 cells with IC50 of 0.3-50 nM, was prepd. in 5 steps involving oxidn. and fractional crystn. using (+)-cinchonine and cinchonidine.

IT 224307-87-3P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); **THU (Therapeutic use)**; BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of optically active 1,4-dihydropyridines as bradykinin antagonists for pharmaceutical compns.)

RN 224307-87-3 HCAPLUS

CN 3,5-Pyridinedicarboxylic acid,

4-(2,6-dichlorophenyl)-1,4-dihydro-2-[2-oxo-

2-[4-[2-(1-pyrrolidinyl)ethyl]-1-piperazinyl]ethyl]-6-[[(S)-phenylsulfinyl]methyl]-, dimethyl ester, dihydrochloride, (4S)- (9CI)

INDEX NAME)

(CA

Absolute stereochemistry. Rotation (-).

L5 ANSWER 2 OF 8 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1997:587253 HCAPLUS

DOCUMENT NUMBER: 127:248125

TITLE: Preparation of 2-(piperazinylcarbonylmethyl)-3,5-

bis(methoxycarbonyl)-1,4-dihydropyridines as

bradykinin antagonists.

INVENTOR(S): Ikeda, Takafumi

PATENT ASSIGNEE(S): Pfizer Pharmaceuticals Inc., Japan; Pfizer Inc.;

Ikeda, Takafumi

SOURCE: PCT Int. Appl., 42 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PAT	TENT	NO.		KI	ND	DATE		APPLICATION NO.					DATE				
WO	9730	048		A	1	1997	0821		M() 19	 97-I	==== В58		1997	0127		
	W:	AU,	BG,	BR,	BY,	CA,	CN,	CZ,	HU,	IL,	IS,	JΡ,	KR,	ΚZ,	LK,	LV,	MX,
		NO,	NΖ,	PL,	RO,	RU,	SG,	SI,	SK,	TR,	UA,	US,	UZ,	VN			
	RW:	AT,	BE,	CH,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙT,	LU,	MC,	NL,	PT,
		SE,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	ML,	MR,	ΝE,	SN,	TD,	TG	
ΑU	9713	964		A	1	1997	0902		ΑI	J 19	97-1	3964		1997	0127		
ΕP	8820	44		Α	1	1998	1209		E	P 19	97-9	0040	1	1997	0127		
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙΤ,	LI,	LU,	NL,	SE,	PT,	ΙE,
		SI,	LV,	FI													
CN	1211	251		Α		1999	0317		CI	N 19	97-1	9224	0	1997	0127		
JΡ	1150	7949		T	2	1999	0713		J!	P 19	97-5	2913	8	1997	0127		
JΡ	3167	335		В	2	2001	0521										
BR	9707	568		Α		1999	0727		B	R 19	97-7	568		1997	0127		
ZA	9701	357		Α		1998	0818		ZI	A 19	97-1	357		1997	0218		

NO 9803776 A 19980818 NO 1998-3776 19980818 US 6131226 A 20001017 US 1999-125137 19990201 PRIORITY APPLN. INFO.: WO 1996-IB132 A 19960219 WO 1997-IB58 W 19970127

OTHER SOURCE(S):

MARPAT 127:248125

GΙ

AB Title compds. [I; A1, A2 = halo; X = C0, SO2, SO(CH2)n; n = 0-2; R1 = 8-azabicyclo[3.2.1]octyl, quinuclidinyl, bicyclo[3.3.0]octyl, cycloalkyl, 2,3,5,6-tetrahydro-4H-thiopyranyl, (substituted) cycloalkylalkyl], were prepd. I have excellent bradykinin antagonistic activity and are useful for the treatment of inflammation, cardiovascular disease, pain, common cold, allergies, asthma, pancreatitis, burns, virus infection, head injury, or multiple trauma. Thus, Me

2-(4,6-dichlorophenylmethylidene)-3-

oxo-4-phenylthiobutanoate (prepn. given) and di-Me 3-aminoglutaconate were

heated at 120.degree. for 3 h to give 40.6% di-Me

4-(2,6-dichlorophenyl)-2-

methoxycarbonylmethyl-6-phenylthiomethyl-1,4-dihydropyridine-3,5-dicarboxylate. This was oxidized to the phenylsulfinyl deriv., which was partially sapond. followed by amidation with 1-(8-methyl-8-azabicyclo[3.2.1]oct-3-yl)piperazine using N-1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride in CH2Cl2 to give di-Me $4-(2,6-\mbox{dichlorophenyl})-2-[4-(8-\mbox{methyl-8-azabicyclo}[3.2.1]oct-3-yl)-1-$

piperazinyl]carbonylmethyl-6-phenylsulfinylmethyl-1,4-dihydropyridine-3,5dicarboxylate dihydrochloride. I inhibited [3H]bradykinin binding to ileum prepns. with IC50 = 0.2-10 nM.

IT 195503-93-6P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); **THU (Therapeutic use)**; BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of 2-(piperazinylcarbonylmethyl)~3,5-bis(methoxycarbonyl)-1,4-dihydropyridines as bradykinin antagonists)

RN 195503-93-6 HCAPLUS

CN 3,5-Pyridinedicarboxylic acid,

4-(2,6-dichlorophenyl)-1,4-dihydro-2-[2-[4-

(8-methyl-8-azabicyclo[3.2.1]oct-3-yl)-1-piperazinyl]-2-oxoethyl]-6-[(phenylsulfinyl)methyl]-, dimethyl ester, dihydrochloride (9CI) (CA INDEX NAME)

● 2 HCl

L5 ANSWER 3 OF 8 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1996:404662 HCAPLUS

DOCUMENT NUMBER: 125:86676

TITLE: Preparation of 2-(piperazinocarbonylmethyl)-1,4-

dihydropyridinedicarboxylates as bradykinin

antagonists

INVENTOR(S): Ito, Fumitaka; Kondo, Hiroshi; Hageman, David L.;

Lowe, John A., III; Nakanishi, Susumu; Vinick,

Fredric

т.

PATENT ASSIGNEE(S): Pfizer Pharmaceuticals Inc., Japan

SOURCE: PCT Int. Appl., 64 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PA:	TENT NO.		KIND	DATE	APPLICATION NO.	DATE
WO	9606083 W: JP		A1	19960229	WO 1994-JP1398	19940824
CA			AA	19960229	CA 1995-2198231	19950526
WO	9606082		A1	19960229	WO 1995-IB400	19950526
	W: AU,	CA,	CN, FI	, JP, KR, MX,	NO, NZ, US	
	RW: AT,	BE,	CH, DE	, DK, ES, FR,	GB, GR, IE, IT, LU	, MC, NL, PT, SE
ΑU				19960314		
ΑU	689587		В2	19980402		
ΕP	777653		A1	19970611	EP 1995-918113	19950526
EΡ	777653		В1	20010822		
	R: AT,	BE,	CH, DE	, DK, ES, FR,	GB, GR, IE, IT, LI	, LU, NL, PT, SE
CN					CN 1995-194758	

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JP 09510992
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                             19971104
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     JP 3001978
                       В2
                             20000124
     AT 204567
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                       Ε
                                             AT 1995-918113
                                                              19950526
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     FI 9700745
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                             19970221
                                             FI 1997-745
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                       Α
                             19990112
                                             US 1997-793561
                                                              19970701
PRIORITY APPLN. INFO.:
                                          WO 1994-JP1398
                                                              19940824
                                          WO 1995-IB400
                                                           W
                                                              19950526
```

OTHER SOURCE(S): MARPAT 125:86676

GΙ

ΙT

AΒ Title compds. [I; R1 = H, (un)substituted (cyclo)alkyl, azacycloalkyl, etc.; R2 = H, alkyl, Ph, etc.; R3,R4 = alkyl; R5,R6 = halo; Z = bond, CH2,

O, CO, etc.] were prepd. Thus, 2,6-Cl2C6H3CH:C(CO2Me)COCH2CH2Ph was cyclocondensed with MeO2CCH:C(NH2)CO2Me (prepn. each given) and the product amidated by N-methylpiperazine to give I (R1 = R3 = R4 = Me, R2 = Ph, R5 = R6 = C1, Z = CH2). I had IC50 of 5nM to 1.mu.M against bradykinin binding at ileum tissue prepn. in vitro. 178376-99-3P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of 2-(piperazinocarbonylmethyl)-1,4-

dihydropyridinedicarboxylates as bradykinin antagonists)

RN 178376-99-3 HCAPLUS

CN 3,5-Pyridinedicarboxylic acid, 4-(2,6-dichlorophenyl)-1,4-dihydro-2-[[(4-

methoxyphenyl)sulfonyl]methyl]-6-[2-(4-methyl-1-piperazinyl)-2-oxoethyl]-, dimethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 4 OF 8 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1991:185276 HCAPLUS

DOCUMENT NUMBER: 114:185276

TITLE: A process for preparation of enantiomerically pure

polysubstituted 1,4-dihydropyridines

INVENTOR(S): Gandolfi, Carmelo A.; Frigerio, Marco; Riva, Carlo;

Zaliani, Andrea; Long, Giorgio; Di Domenico, Roberto

PATENT ASSIGNEE(S): Boehringer Biochemia Robin S.p.A., Italy

SOURCE: Eur. Pat. Appl., 32 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PA'	rent 	NO.	KII	ND	DATE		Al	PPLI	CATIO	ON NO	Ο.	DATE			
EP	3833 R:		A	1	1990	0822	El	2 19	90-1	0295	1	1990	0215		
	2047 9009														
		ΑU,	BG,									MG,		NO,	RO,
	RW:				CF, SN,		CM,	DE,	DK,	ES,	FR,	GA,	GB,	IT,	LU,
	9050 6309						Ą	J 19	90-50	0904		1990	0215		
	4588 4588						El	2 19:	90-90	03019	5	1990	0215		
HU AT	R: 0450 6227 9581 2060	0	T: A:	2 2	1992: 1993:	1001 0428 1015	JI HI A:	P 19: J 19: Γ 19:	90-50 90-19 90-90	03175 962 03015	5	SE 19900 19900 19900	0215 0215		

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RU 1990-5001680 19900215
     RU 2069658
                        C1
                             19961127
     US 5245039
                                                                19910814
                              19930914
                                              US 1991-743415
                        Α
     NO 9103188
                              19910815
                                              NO 1991-3188
                        Α
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                        В
                              19950424
     NO 177186
                        С
     NO 177186
                              19950802
                        В
     FI 95371
                              19951013
                                              FI 1991-3861
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     FI 95371
                        С
                              19960125
PRIORITY APPLN. INFO.:
                                           IT 1989-19477
                                                                19890217
                                          EP 1990-903015
                                                                19900215
                                          WO 1990-EP243
                                                                19900215
OTHER SOURCE(S):
                         MARPAT 114:185276
GΙ
                                       NO2
      R4
           R3
                         Et02C
                                        CO<sub>2</sub>Et
                                                    ΙI
AΒ
     The title compds. [I; R3 = (esterified) CO2H; R4 = (substituted) Ph,
     .beta.-naphthyl, heterocyclyl, etc.; R5 = cyano, NO2, (esterified) CO2H,
     etc.; R6 = C1-6 alkyl halo-C1-6-alkyl, HOC, CN, etc.; A = H,
     isothioureido, SH, sulfonium salt, etc.; n = 1-4], useful as cardiovascular agents (no data), are prepd. A mixt. of 6 g (.+-.)-I (A =
     C1, R3 = R5 = C02Et, R4 = 3-02NC6H4, R6 = Me, n = 1) and 1.2 g thiourea
in
     EtOH was refluxed to give 4.8 g isothiuronium salt (.+-.)-II.HCl, which
     was treated with NaHCO2 in EtOAc-H2O to give free (.+-.)II. Optical
     resoln. of (.+-.)-II with O,O'-dibenzoyl-D-tartaric acid gave (+)-II of
     >98% optical purity. Also prepd. were over 100 chiral I.
ΙT
     131767-73-2P
     RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study);
     PREP (Preparation); USES (Uses)
        (prepn. of, as cardiovascular agent)
     131767-73-2 HCAPLUS
RN
CN
     Sulfoxonium,
[[3-(ethoxycarbonyl)-1,4-dihydro-5-(methoxycarbonyl)-6-methyl-
     4-(3-nitrophenyl)-2-pyridinyl]methyl]dimethyl-, (+)-,
tetrafluoroborate(1-
     ) (9CI) (CA INDEX NAME)
     CM
          1
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CRN 131767-72-1 CMF C20 H25 N2 O7 S

CDES 3: (+)

Rotation (+).

CM 2

CRN 14874-70-5

CMF B F4

L5 ANSWER 5 OF 8 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: 1990:515098 HCAPLUS

DOCUMENT NUMBER: 113:115098
TITLE: Preparation of 2-thiomethylpyridine-3,5-dicarboxylates

as antihypertensives

INVENTOR(S): Gandolfi, Carmelo A.; Frigerio, Marco; Tofanetti,

Odoardo; Tognella, Sergio

PATENT ASSIGNEE(S): Boehringer Biochemia Robin S.p.A., Italy

SOURCE: U.S., 8 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

KIND DATE PATENT NO. APPLICATION NO. DATE --------------A US 1988-167164 JP 1988-58141 US 4918087 19900417 19880311 A2 JP 63243073 19881007 19880311 PRIORITY APPLN. INFO.: IT 1987-19700 19870313 OTHER SOURCE(S): MARPAT 113:115098

$$R^{2}$$
 R^{1}
 R^{3}
 R^{3}
 R^{3}
 R^{2}
 R^{3}
 R^{3}
 R^{3}
 R^{4}
 R^{3}

AB The title compds. (I; R1, R3 = CO2R5; R2 = m-O2NC6H4, m-NCC6H4; R4 = (CH2)mCH2NR6R7; R5 = C1-6 alkyl; R6, R7 = H; 1 of R6, R7 = H, the other

C1-6 alkyl, PhCH2) and their pharmaceutically acceptable salts, antihypertensives without Ca2+-antagonizing activity, were prepd. by aromatization of their 1,4-dihydropyridine analogs or by thiolation of 2-halomethylpyridine precursors. Thus, a soln. of 8.5 g 2-chloromethyl-3-carboethoxy-5-carbomethoxy-4-(m-nitrophenyl)-6-methylpyridine in ethane was added dropwise to a soln. of 1.6 g H2NCH2CH2SH.HCl and 7.3 mL 20% NaOH in 50 mL EtOH at -10.degree.. After 15 min the soln. was warmed up to room temp. to give 9 g I (R1 = CO2Me,

R2 = m-O2NC6H4, R3 = CO2Et, R4 = CH2CH2NH2, n = 0) which was converted to its

fumarate salt. In spontaneously hypertensive rats, the latter (form unspecified) at 3 mg/kg orally reduced mean blood pressure by 10-20%.

118587-28-3P
RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of, as antihypertensive)

RN 118587-28-3 HCAPLUS

ΙT

CN 3,5-Pyridinedicarboxylic acid,

2-[[(2-aminoethyl)sulfinyl]methyl]-6-methyl-

4-(3-nitrophenyl)-, 3-ethyl 5-methyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 6 OF 8 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1988:610906 HCAPLUS

DOCUMENT NUMBER: 109:210906

TITLE: 2-Methylthiomethyldihydropyridines, a process for

their preparation, and pharmaceutical compositions containing them useful as cardiovascular agents Frigerio, Marco; Zaliani, Andrea; Riva, Carlo;

Gandolfi, Carmelo A.; Tofanetti, Odoardo; Tognella,

Sergio

PATENT ASSIGNEE(S): Boehringer Biochemia Robin S.p.A., Italy

SOURCE: Eur. Pat. Appl., 38 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

INVENTOR(S):

PA'	TENT NO.	K	IND I	DATE		APPLICATION NO.	DATE
EP	271059		A1 :	19880615		EP 1987-118145	19871208
EP	271059		В1 :	19931020			
	R: AT,	BE, CH	, DE,	ES, FR,	GB, GI	R, IT, LI, LU, NL	, SE
AT	96152		Ε :	19931115		AT 1987-118145	19871208
AU	8782294	i	A1 :	19880616		AU 1987-82294	19871210
AU	611872		B2 :	19910627			
JP	63170360		42	19880714		JP 1987-313104	19871210
ZA	8709301		Α .	19880831		ZA 1987-9301	19871210
US	4971984		Α :	19901120		US 1987-131393	19871210
PRIORIT'	Y APPLN.	INFO.:			ΙT	1986-22648	19861211
					EΡ	1987-118145	19871208

OTHER SOURCE(S): MARPAT 109:210906

GΙ

$$R^{1}$$
 R^{2}
 R^{3}
 R^{3}
 R^{2}
 R^{3}
 R^{3}
 R^{2}
 R^{3}
 R^{3}
 R^{2}
 R^{3}
 R^{3

AB Title compds. I [R1 = Ac, Bz, cyano, O2N, R7O2C; R7 = (un)substituted C1-6

alkyl, C3-6 alkenyl, (un)substituted Ph, R8R9NCO; R8, R9 = H, C1-6 alkyl,
PhCH2, aryl; R2 = (un)substituted Ph, F5C6, .alpha.-, .beta.-naphthyl,
5-6

membered heterocyclyl; R3 = R7O2C; R4, R5 = H, C1-6 alkyl, (un) substituted

Ph, R7O2C, C3-7 cycloalkyl, 5-6 membered heterocyclyl; R4R5C = 5-6 membered ring; Y = 0, S, R10N:; R6, R10 = H, (un)substituted C1-6 alkyl, C1-12 alkanoyl, (un)substituted aroyl or heteroaryl, (un)substituted aryl or 5-6 membered heteroaryl contg. O, S, N; R6R10N = pyrrolyl, piperidyl, morpholyl, piperazinyl, succinimidyl, phthaloyl; n = 0-2] useful as cardiovascular agents (no data) were prepd.

3,5-Dicarboethoxy-6-methyl-2-

(mercaptomethyl)-4-(m-nitrophenyl)-1,4-dihydropyridine, and sulfosalicylic

acid in MeC(OMe)2Me were stirred for 2 h at room temp. to give I (R1 = \pm tO2C; R2 = 4-(O2N)C6H4; R3 = CO2Et; R4, R5, R6 = Me; Y = O; n = 0).

IT 116508-72-6P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); **THU (Therapeutic use)**; BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of, as cardiovascular agent)

RN 116508-72-6 HCAPLUS

CN 3,5-Pyridinedicarboxylic acid,

2-[[[(acetyloxy)methyl]sulfinyl]methyl]-1,4-

dihydro-6-methyl-4-(3-nitrophenyl)-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 7 OF 8 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1988:21721 HCAPLUS

DOCUMENT NUMBER: 108:21721

TITLE: Preparation of dihydro(thiomethyl)pyridines as

antihypertensives

INVENTOR(S): Gandolfi, A. Carmelo; Frigerio, Marco; Spinelli,

Silvano; Tofanetti, Odoardo; Tognella, Sergio

PATENT ASSIGNEE(S): Boehringer Biochemia Robin S.p.A., Italy

SOURCE: PCT Int. Appl., 91 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PAT	TENT	NO.		KI	ND	DATE			A	PPLI	CATI	ON NO	Ο.	DATE			
WO	8700	836		A	1	1987	0212		M) 19	86-E	P445		1986	729		
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		SD,	SU														
	RW:	ΑT,	BE,	CF,	CG,	CH,	CM,	DE,	FR,	GΑ,	GB,	ΙΤ,	LU,	ML,	MR,	NL,	SE,
		SN,	TD,	ΤG													
ΑU	8662	245		Α	1	1987	0305		A	J 19	86-6	2245		1986	0729		
ΑU	5932	78		В	2	1990	0208										
EΡ	2332	28		А	1	1987	0826		E	P 19	86-9	0481	3	1986	0729		
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HU	4449	9		A	2	1988	0328		H	J 19	86-4	090		1986	729		
НU	1997	96		В		1990	0328										
IL	7957	2		Α	1	1991	0630		I	L 19	86-7	9572		1986	0730		
CN	8610	6242		А		1987	0513		C	N 19	86-1	0624	2	1986	0805		

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PRIORITY APPLN. INFO.:
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                                                          19870625
                                       WO 1987-EP335
                                                          19870625
                                       US 1989-299992
                                                          19890221
GΙ
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The title compds. [I; R1 = acyl, cyano, NO2, alkoxycarbonyl, H2NCO; R2 = (un)substituted aryl, heteroaryl; R3 = CO2R4; R4 = H, alkenyl, (un)substituted alkyl, aryl, aralkyl; X = S(O)nR5; R5 = H, acyl, (un)substituted alkyl, alkenyl, alkynyl, cycloalkyl, aryl, heterocyclyl, alkoxyalkyl, alkylthioalkyl, thiuronium; n = 0-2] were prepd. as antihypertensives. Dihydropyridinedicarboxylate II (Y = Cl) (6 g) was refluxed with 1.2 g (H2N)2CS in EtOH to give 4.8 g dihydropyridinyisothiuronium salt II.HCl [Y = SC(:NH)NH2] (III). III had an IC50 of 2.2 .times. 10-7 M for inhibition of aortal-strip contraction and in rats 3.1 mg III/kg orally reduced blood pressure 45 mmHg.

IT 110645-87-9P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); **THU (Therapeutic use)**; BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of, as antihypertensive)

RN 110645-87-9 HCAPLUS

CN 3,5-Pyridinedicarboxylic acid, 1,4-dihydro-2-methyl-4-(3-nitrophenyl)-6-[(phenylsulfinyl)methyl]-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 8 OF 8 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1987:176173 HCAPLUS

DOCUMENT NUMBER: 106:176173

TITLE: 1,4-Dihydropyridine derivatives useful in the

treatment of cardiovascular disorders

INVENTOR(S):
Sircar, Ila

PATENT ASSIGNEE(S): Warner-Lambert Co., USA SOURCE: Eur. Pat. Appl., 80 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO. DATE	
EP 206747 EP 206747	A2 A3	19861230 19870819	EP 1986-304679 198606	17
	B1 CH, DE	19901031 , FR, GB,	IT, LI, LU, NL, SE	
ZA 8603914	А	19880127	ZA 1986-3914 198605	26
FI 8602343	Α	19861218	FI 1986-2343 198606	02
CA 1267416	A1	19900403	CA 1986-510582 198606	02
AU 8658428	A1	19861224	AU 1986-58428 198606	04
AU 601946	В2	19900927		
DK 8602811	A	19861218	DK 1986-2811 198606	16
NO 8602390	А	19861218	NO 1986-2390 198606	16
JP 62036357	A2	19870217	JP 1986-138417 198606	16
ES 556082	A1	19871201	ES 1986-556082 198606	16
CN 86104284	A	19870401	CN 1986-104284 198606	17
AT 57915	E	19901115	AT 1986-304679 198606	17
PRIORITY APPLN. INFO.	:		US 1985-745965 198506	17
			US 1986-852731 198604	21
			EP 1986-304679 198606	17
GI				

 $(CH_2)_{m}S(0)_{n}R^{1}$

AB The title compds. [I; R = H, C1-4 alkyl, optionally substituted with R7R8N, (un)substituted aralkyl; R1, R5, R6 = H, (un)substituted C1-4 alkyl; R2, R3 = H, C1-4 alkyl, cyano, NO2, CO2R4; R2R5 = atoms to complete

a 5- or 6-membered carbocycle; R4 = H, (un)substituted C1-4 alkyl; R7, R8 = H, alkyl; R7R8N = 5- or 6-membered ring, m = 1-6; n = 0-2] and their salts, having Ca antagonist properties (no data) and inotropic properties,

were prepd. I can be formulated into pharmaceuticals (no data). Thus, 2-F3CC6H4CHO, H2NCMe:CHCN, and PhSCH2COCH2CO2Et in EtOH were refluxed to give I (R = H, R1 = Ph, R3 = CO2Et, R5 = Me, R6 = 2-F3CC6H4, m = 1, n = $\frac{1}{2}$

double bond present) which was oxidized to give I (n=1, other variables the same). Myocardial inotropic activity in isolated guinea pig atria was

demonstrated with selected I.

IT 107975-19-9P

0,

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); **THU (Therapeutic use)**; BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of, as cardiovascular agent)

107975-19-9 HCAPLUS RN

CN 3,5-Pyridinedicarboxylic acid, 1,4-dihydro-2-methyl-6-[(4pyridinylsulfinyl)methyl]-4-[2-(trifluoromethyl)phenyl]-, diethyl ester (9CI) (CA INDEX NAME)

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FULL ESTIMATED COST	ENTRY 51.74	SESSION 192.43
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This file contains CAS Registry Numbers for easy and accurate substance identification. Title keywords, authors, patent assignees, and patent information, e.g., patent numbers, are now searchable from 1907-1966. TIFF images of CA abstracts printed between 1907-1966 are available in the PAGE display formats.

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